Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	4533	cell near9 pore	US-PGPUB; USPAT; EPO; DERWENT	OR	OFF	2005/01/24 13:28
L2	О	I1 same electric?	US-PGPUB; USPAT; EPO; DERWENT	OR	OFF	2005/01/24 13:28
L3	408	I1 same (electrical, potential, capacitance, resistance, conductance)	US-PGPUB; USPAT; EPO; DERWENT	OR	OFF	2005/01/24 13:29
L4	104	I3 same surface	US-PGPUB; USPAT; EPO; DERWENT	OR	OFF	2005/01/24 13:29
L5	84	l4 and @py<"2004"	US-PGPUB; USPAT;	OR	OFF	2005/01/24 13:45
			EPO; DERWENT			
L6	0	("9886360").PN.	USPAT; EPO	OR	OFF	2005/01/24 13:46
L7	1	("6767515").PN.	USPAT; EPO	OR	OFF	2005/01/24 13:46

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FULL ESTIMATED COST

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=> pore and cell and electrical
L1 24 FILE AGRICOLA
L2 118 FILE BIOTECHNO
L3 0 FILE CONFSCI
L4 1 FILE HEALSAFE
L5 0 FILE IMSDRUGCONF
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L6 107 FILE LIFESCI
L7 0 FILE MEDICONF

L8 250 FILE PASCAL

TOTAL FOR ALL FILES

500 PORE AND CELL AND ELECTRICAL

=> 19 and (surface or top)
L10 1 FILE AGRICOLA

L11 21 FILE BIOTECHNO L12 0 FILE CONFSCI

L13 0 FILE HEALSAFE

L14 0 FILE IMSDRUGCONF

L15 15 FILE LIFESCI L16 0 FILE MEDICONF

L17 78 FILE PASCAL

TOTAL FOR ALL FILES

L18 115 L9 AND (SURFACE OR TOP)

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L19
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L20
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             0 FILE CONFSCI
L21
             O FILE HEALSAFE
L22
L23
             0 FILE IMSDRUGCONF
             O FILE LIFESCI
L24
             O FILE MEDICONF
L25
L26
             0 FILE PASCAL
TOTAL FOR ALL FILES
             0 L9 AND CELL ATTACHMENT
L27
=> 19 and attachment
             0 FILE AGRICOLA
             0 FILE BIOTECHNO
L29
             O FILE CONFSCI
L30
             O FILE HEALSAFE
L31
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L32
             O FILE LIFESCI
L33
L34
             O FILE MEDICONF
             2 FILE PASCAL
L35
TOTAL FOR ALL FILES
             2 L9 AND ATTACHMENT
=> d 136 ibib abs total
      ANSWER 1 OF 2 PASCAL COPYRIGHT 2005 INIST-CNRS. ALL RIGHTS RESERVED. on
      STN
                                         PASCAL
ACCESSION NUMBER:
                         1997-0476832
                         Copyright .COPYRGT. 1997 INIST-CNRS. All rights
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                         reserved.
                         Pulsed-laser metal contacting of biosensors on the
TITLE (IN ENGLISH):
                         basis of crystalline enzyme-protein layer composites
                         NEUBAUER A.; PENTZIEN S.; REETZ S.; KAUTEK W.; PUM D.;
AUTHOR:
                         SLEYTR U. B.
                         Laboratory for Thin Film Technology, Federal Institute
CORPORATE SOURCE:
                         for Materals Research and Testing, Unter den Eichen
                         87, 12205 Berlin, Germany, Federal Republic of;
                         Nanosearch Membrane Ges.m.b.H. (NSM), Hettenkofergasse
                         13/45, 1160 Vienna, Austria; Center for Ultrastructure
                         Research, Vienna University of Agriculture, Forestry
                         and Renewable Natural Resources, Gregor-Mendel-Strasse
                         33, 1180 Vienna, Austria
                         Sensors and actuators. B, Chemical, (1997), 40(2-3),
SOURCE:
                         231-236, 22 refs.
                         ISSN: 0925-4005
DOCUMENT TYPE:
                         Journal
BIBLIOGRAPHIC LEVEL:
                         Analytic
COUNTRY:
                         Switzerland
LANGUAGE:
                         English
                         INIST-19425B, 354000068856110230
AVAILABILITY:
                     PASCAL
      1997-0476832
AN
      Copyright .COPYRGT. 1997 INIST-CNRS. All rights reserved.
CP
      Crystalline bacterial cell surface layers (S-layers) composed
AΒ
      of monomolecular arrays of protein subunits are accessible to-a wide
      variety of possible proteinchemical reactions. This enables the
      attachment and immobilization of enzyme molecules-in a tightest
      packing, which has not been achieved with other immobilization matrices.
      When immobilized to an S-layer lattice, the enzyme entities are
      surrounded by nanometer pores. Thus, they can react
      electrochemically with the analyte liquid streaming through these
```

=> 19 and cell attachment

pores. The control over this process has to take place by way of an inert electrical contact in a distance of less than 1 nm. The relatively voluminous, but specially shaped sensor enzyme molecules have to be connected with an optimum metallic contact, which must not disturb the protein structure. Previously, platinum films were applied on enzyme layers immobilized on S-layer protein by argon sputtering. This conventional technique, however, exhibits substantial limitations. One, for instance, is the volume change of the S-layer/enzyme composite system when it is introduced into a conventional vacuum coating apparatus. This coating problem can be circumvented by a completely new deposition method, i.e. the pulse-laser-deposition (PLD) on protein crystal composite films with optimized laser parameters and reaction atmospheres. Enzyme activities of 70-80% were achieved, thus demonstrating that composite systems consisting of the 2D-protein-layer/enzyme/metal sequence can successfully serve as highly efficient sensor systems.

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STN

ACCESSION NUMBER: 1996-0190415 PASCAL

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reserved.

TITLE (IN ENGLISH): Enhanced transport of bacteria in porous media by

sediment-phase and aqueous-phase natural organic

matter

AUTHOR: JOHNSON W. P.; LOGAN B. E.

CORPORATE SOURCE: Department of Geology and Geophysics, The University

of Utah, Salt Lake City, UT 84112, United States Water research: (Oxford), (1996), 30(4), 923-931, 3

tabl., refs. 1 p. ¬ Illustrations; Table

ISSN: 0043-1354 CODEN: WATRAG

DOCUMENT TYPE:

SOURCE:

BIBLIOGRAPHIC LEVEL: Analytic

COUNTRY: United Kingdom

LANGUAGE: English

AVAILABILITY: INIST-8940A, 354000044829730190

Journal

AN 1996-0190415 PASCAL

CP Copyright .COPYRGT. 1996 INIST-CNRS. All rights reserved.

Aqueous-phase dissolved natural organic matter (DOM) and sediment organic AR matter (SOM) were shown in laboratory mini-column experiments to affect the transport of bacteria within porous media. Attachment efficiencies of bacteria were estimated from their retention on quartz, iron oxide coated quartz (Fe-quartz), and Fe-quartz coated with SOM (SOM-Fe-quartz). Suwannee River Humic Acid (SRHA) and Soil Humic Acid (SHA) were used to represent organic matter (SOM and DOM), and were added to radiolabeled bacterial suspensions (10.sup.6 cells/ml, pH = 7.7) prior to transport. Coating quartz with iron oxide increased bacterial retention 160% relative to uncoated quartz. Coating Fe-quartz with SOM lowered bacterial retention, resulting in a fraction retained only 33% greater than retained on uncoated quartz. Compared to these effects, the effect of DOM on bacterial retention was secondary, and reflected the extent of DOM adsorption to the porous media. When DOM did not interact with the porous media, as in the case of quartz, bacterial retention in the presence of DOM was reduced by 20%. However, when DOM adsorption to the porous media was increased by coating the quartz with iron oxide, bacterial retention on the Fe-quartz increased by 10%. When Fe-quartz surfaces were loaded with DOM to equilibrium conditions to produce SOM-Fe-quartz, the presence of DOM in the applied solution also increased bacterial retention by 10%. The effects of DOM were the same for both types of humic acids (SHA or SRHA). These results suggest that SOM and DOM affect bacterial transport by increasing the negative surface charge of the Fe-quartz and bacteria, respectively. The largest decrease in bacterial retention (60%) was associated with coating of Fe-quartz by SOM in the absence of DOM.

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11 FILE AGRICOLA
L37
L38
            57 FILE BIOTECHNO
L39
            0 FILE CONFSCI
            O FILE HEALSAFE
L40
            O FILE IMSDRUGCONF
L41
            28 FILE LIFESCI
L42
            O FILE MEDICONF
L43
            56 FILE PASCAL
L44
TOTAL FOR ALL FILES
          152 L9 AND POTENTIAL
L45
=> 19 and action potential
            O FILE AGRICOLA
L46
            10 FILE BIOTECHNO
L47
             0 FILE CONFSCI
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             O FILE HEALSAFE
L49
             O FILE IMSDRUGCONF
L50
             6 FILE LIFESCI
L51
             0 FILE MEDICONF
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             8 FILE PASCAL
L53
TOTAL FOR ALL FILES
            24 L9 AND ACTION POTENTIAL
L54
=> dup rem
ENTER L# LIST OR (END):154
DUPLICATE IS NOT AVAILABLE IN 'IMSDRUGCONF, MEDICONF'.
ANSWERS FROM THESE FILES WILL BE CONSIDERED UNIQUE
PROCESSING COMPLETED FOR L54
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=> d 155 ibib abs total
      ANSWER 1 OF 19 PASCAL COPYRIGHT 2005 INIST-CNRS. ALL RIGHTS RESERVED.
      on STN
ACCESSION NUMBER:
                         2004-0298608
                                         PASCAL
                         Copyright .COPYRGT. 2004 INIST-CNRS. All rights
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                         Suppression of electrical alternans by
TITLE (IN ENGLISH):
                         overexpression of HERG in canine ventricular myocytes
                         FEI HUA; JOHNS David C.; GILMOUR Robert F. JR
AUTHOR:
                         Department of Biomedical Sciences, Cornell University,
CORPORATE SOURCE:
                         Ithaca, New York 14853, United States; Department of
                         Neurosurgery, Johns Hopkins School of Medicine,
                         Baltimore, Maryland 21205, United States
                         American journal of physiology. Heart and circulatory
SOURCE:
                         physiology, (2004), 55(6), 2342-2352, 39 refs.
                         ISSN: 0363-6135 CODEN: AJPPDI
                         Journal
DOCUMENT TYPE:
                         Analytic
BIBLIOGRAPHIC LEVEL:
                         United States
COUNTRY:
LANGUAGE:
                         English
                         INIST-670D, 354000111978950380
AVAILABILITY:
      2004-0298608
                     PASCAL
AN
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CP
      Suppression of electrical alternans may be antiarrhythmic. Our
AB
      previous computer simulations have suggested that increasing the rapid
      component of the delayed rectifier K.sup.+ current (I.sub.K.sub.r)
```

suppresses alternans. To test this hypothesis, I.sub.K.sub.r in isolated canine ventricular myocytes was increased by infection with an adenovirus

=> 19 and potential

containing the gene for the pore-forming domain of I.sub.K.sub.r [human ether-a-go-go gene (HERG)]. With the use of the perforated or whole cell patch-clamp technique, action potentials recorded at different pacing cycle lengths (CLs) were applied to the myocytes as the command waveforms. HERG infection markedly increased peak I.sub.K.sub.r during the action potential (from 0.54 \pm 0.03 pA/pF in control to 3.60 \pm 0.81 pA/pF). Rate-dependent alterations of peak I.sub.K.sub.r were similar for freshly isolated myocytes and HERG-infected myocytes. In both cell types, I.sub.K.sub.r increased when CL decreased from 1,000 to 500 ms and then decreased progressively as CL decreased further. During alternans at CL = 170 ms, peak I.sub.K.sub.r was larger for the short than for the long action potential for both groups, but the difference in peak I.sub.K.sub.r was larger for HERG-infected myocytes. The voltage at which peak I.sub.K.sub.r occurred was significantly less negative in HERG-infected myocytes, in association with shifts of the steady-state voltage-dependent activation and inactivation curves to less negative potentials. Pacing at short CL induced stable alternans in freshly isolated myocytes and in cultured myocytes without HERG infection, but not in HERG-infected myocytes. These data support the idea that increasing I.sub.K.sub.r may be a viable approach to suppressing electrical alternans.

ANSWER 2 OF 19 PASCAL COPYRIGHT 2005 INIST-CNRS. ALL RIGHTS RESERVED. L55

on STN

ACCESSION NUMBER:

2004-0594545 PASCAL

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TITLE (IN ENGLISH):

Heart rate lowering by specific and selective If current inhibition with ivabradine: A new therapeutic

perspective in cardiovascular disease

AUTHOR:

SOURCE:

AB

DIFRANCESCO Dario; CAMM John A.

CORPORATE SOURCE:

Dipartimento di Scienze Biomolecolari e Biotecnologie,

Universita di Milano, Milan, Italy; The Medical

School, St George's Hospital, London, United Kingdom Drugs: (Basel), (2004), 64(16), 1757-1765, 66 refs. ISSN: 0012-6667 CODEN: DRUGAY

Journal DOCUMENT TYPE: Analytic BIBLIOGRAPHIC LEVEL: COUNTRY: New Zealand English

LANGUAGE: AVAILABILITY:

INIST-15326, 354000122211320030

2004-0594545 PASCAL AN

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Resting heart rate is associated with cardiovascular and all-cause mortality, and the mortality benefit of some cardiovascular drugs seems to be related in part to their heart rate-lowering effects. Since it is difficult to separate the benefit of heart rate lowering from other actions with currently available drugs, a 'pure' heart rate-lowering drug would be of great interest in establishing the benefit of heart rate reduction per se. Heart rate is determined by spontaneous electrical pacemaker activity in the sinoatrial node. Cardiac pacemaker cells generate the spontaneous slow diastolic depolarisation that drives the membrane voltage away from a hyperpolarised level towards the threshold level for initiating a subsequent action potential, generating rhythmic action potentials that propagate through the heart and trigger myocardial contraction. The If current is an ionic current that determines the slope of the diastolic depolarisation, which in turn controls the heart beating rate. Ivabradine is the first specific heart rate-lowering agent to have completed clinical development for stable angina pectoris. Ivabradine specifically blocks cardiac pacemaker cell f-channels by entering and binding to a site in the channel pore from the intracellular side. Ivabradine is selective for the

I.sub.f current and exerts significant inhibition of this current and heart rate reduction at concentrations that do not affect other cardiac ionic currents. This activity translates into specific heart rate reduction, which reduces myocardial oxygen demand and simultaneously improves oxygen supply, by prolonging diastole and thus allowing increased coronary flow and myocardial perfusion. Ivabradine lowers heart rate without any negative inotropic or lusitropic effect, thus preserving ventricular contractility. Ivabradine was shown to reduce resting heart rate without modifying any major electrophysiological parameters not related to heart rate. In patients with left ventricular dysfunction, ivabradine reduced resting heart rate without altering myocardial contractility. Thus, pure heart rate lowering can be achieved in the clinic as a result of specific and selective I.sub.f current inhibition. Two randomised clinical studies have shown that ivabradine is an effective anti-ischaemic agent that reduces heart rate and improves exercise capacity in patients with stable angina. Ivabradine was shown to be superior to placebo in improving exercise tolerance test (ETT) criteria (n = 360) and, in a 4-month, double-blind, controlled study (n = 939), ivabradine 5 and 7.5mg twice daily were shown to be at least as effective as atenolol 50 and 100mg once daily, respectively, in improving total exercise duration and other ETT criteria, and reducing the number of angina attacks. Experimental data indicate a potential role of pure heart rate lowering in other cardiovascular conditions, such as heart failure.

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on STN

ACCESSION NUMBER: 2005-0016624 PASCAL

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TITLE (IN ENGLISH): Transgenic upregulation of I.sub.K.sub.1 in the- mouse

heart leads to multiple abnormalities of cardiac

rearc reads to murciple abnormalities .

excitability

AUTHOR: JINGDONG LI; MCLERIE Meredith; LOPATIN Anatoli N.

CORPORATE SOURCE: Department of Molecular and Integrative Physiology,

University of Michigan Medical School, Ann Arbor,

Michigan 48109, United States

SOURCE: American journal of physiology. Heart and circulatory

physiology, (2004), 56(6), H2790-H2802, 42 refs.

ISSN: 0363-6135 CODEN: AJPPDI

DOCUMENT TYPE:

AB

BIBLIOGRAPHIC LEVEL: Analytic

COUNTRY: United States

LANGUAGE: English

AVAILABILITY: INIST-670D, 354000122548360520

Journal

AN 2005-0016624 PASCAL

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To assess the functional significanc of upregulation of the cardiac current (I.sub.K.sub.1), we have produced anc characterized the first transgenic (TG) mouse model of I.sub.K.sub.1 upregula tion. To increase I.sub.K.sub.1 density, a pore-forming subunit of the Kir2.1 (green fluorescent protein-tagged) channel was expressed in the hear under control of the α -myosin heavy chain promoter. Two lines of TC animals were established with a high level of TG expression in al major parts of the heart: line 1 mice were characterized by 14% hear hypertrophy and a normal life span; line 2 mice displayed an increase mortality rate, and in mice <=1 mo old, heart weight-to-body weigh ratio was increased by >100%. In adult ventricular myocytes expressing the Kir2.1-GFP subunit, I.sub.K.sub.1 conductance at the reversal potentia was increased .eqvsim.9- and .eqvsim.10-fold in lines I and 2, respectively Expression of the Kir2. transgene in line 2 ventricular myocytes was heterogeneous when assayed by single-cell analysis of GFP fluores cence. Surface ECG recordings in line 2 mice revealed numerous abnormalities of excitability, including slowed heart rate,

premature ventricular contractions, atrioventricular block, and atrial fibrillation Line 1 mice displayed a less severe phenotype. In both TG lines action potential duration at 90% repolarization and monophasic actior potential at 75-90% repolarization were significantly reduced, leading to neuronlike action potentials, and the slow phase of the T wave way abolished, leading to a short Q-T interval. This study provides a new TG model of I.sub.K.sub.1 upregulation, confirms the significant role of I.sub.K.sub.lir cardiac excitability, and is consistent with adverse effects of I.sub.K upregulation on cardiac electrical activity.

ANSWER 4 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN L55

BIOTECHNO 2003:36183358 ACCESSION NUMBER:

It takes two to tango, but three to I.sub.S.sub.A TITLE:

Herson P.S.; Adelman J.P. **AUTHOR:**

P.S. Herson, Vollum Institute, Oregon Hlth. and CORPORATE SOURCE:

Sciences University, 3181 S.W. Sam Jackson Park Road,

Portland, OR 97239, United States.

Neuron, (06 FEB 2003), 37/3 (3.70-372), 6 reference(s) SOURCE:

CODEN: NERNET ISSN: 0896-6273

Journal; (Short Survey) DOCUMENT TYPE:

United States COUNTRY:

LANGUAGE: English English SUMMARY LANGUAGE: BIOTECHNO 2003:36183358

cloned subunits.

Rapidly inactivating A-type potassium channels are important determinants AB of firing frequency in many excitable cells. Nadal et al. (in this issue of Neuron) purified A-type potassium (I.sub.S.sub.A) channels from rat cerebellum and identified a novel β subunit. This protein, DPPX, associates with the pore-forming subunits and endows previously elusive kinetic properties on A-type channels formed from

ANSWER 5 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN

2002:34252231 BIOTECHNO ACCESSION NUMBER:

An unexpected role for brain-type sodium channels in TITLE:

coupling of cell surface depolarization to

contraction in the heart

Maier S.K.G.; Westenbroek R.E.; Schenkman K.A.; Feigl AUTHOR:

E.O.; Scheuer T.; Catterall W.A.

W.A. Catterall, University of Washington, Department CORPORATE SOURCE:

of Pharmacology, Campus Box 357280, 1959 NE Pacific Street, Seattle, WA 98195, United States.

E-mail: wcatt@u.washington.edu

Proceedings of the National Academy of Sciences of the SOURCE:

United States of America, (19 MAR 2002), 99/6

(4073-4078), 33 reference(s) CODEN: PNASA6 ISSN: 0027-8424

Journal; Article DOCUMENT TYPE:

United States COUNTRY:

English LANGUAGE: SUMMARY LANGUAGE: English BIOTECHNO AN 2002:34252231

Voltage-gated sodium channels composed of pore-forming α AB and auxiliary β subunits are responsible for the rising phase of the action potential in cardiac muscle, but the functional roles of distinct sodium channel subtypes have not been clearly defined. Immunocytochemical studies show that the principal cardiac poreforming α subunit isoform Na.sub.v1.5 is preferentially localized in intercalated disks, whereas the brain $\boldsymbol{\alpha}$ subunit isoforms Na.sub.v1.1, Na.sub.v1.3, and Na.sub.v1.6 are localized in the transverse tubules. Sodium currents due to the highly tetrodotoxin (TTX)-sensitive brain isoforms in the transverse tubules are small and are detectable only after activation with $\boldsymbol{\beta}$ scorpion toxin. Nevertheless, they play

an important role in coupling depolarization of the cell surface membrane to contraction, because low TTX concentrations reduce left ventricular function. Our results suggest that the principal cardiac isoform in the intercalated disks is primarily responsible for action potential conduction between cells and reveal an unexpected role for brain sodium channel isoforms in the transverse tubules in coupling electrical excitation to contraction in cardiac muscle.

ANSWER 6 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN

2002:34625487 BIOTECHNO ACCESSION NUMBER:

New perspectives on the structure and function of the TITLE:

Na.sup.+ channel multigene family

Ogata N.; Yoshida S. AUTHOR:

N. Ogata, Department of Physiology, Hiroshima Univ. CORPORATE SOURCE:

Sch. of Medicine, Hiroshima 734-8551, Japan.

E-mail: ogatan@hiroshima-u.ac.jp

Current Medicinal Chemistry - Central Nervous System SOURCE:

Agents, (2002), 2/1 (59-81), 108 reference(s)

CODEN: CMCCCO ISSN: 1568-0150

DOCUMENT TYPE:

Journal; General Review

COUNTRY: LANGUAGE:

AN

AB

Netherlands

English SUMMARY LANGUAGE: English BIOTECHNO 2002:34625487

Recent studies on the voltage-gated Na.sup.+ channel (VGSC) have revealed several excellent discoveries regarding its structure and function. This article summarizes recent findings on VGSCs, and presents our views on the subject. Based on the multi-pore 3D model of the VGSC, we propose a "twist-sprinkler" model: (i) twisting and untwisting of the central cavity corresponds to the closed and open states of the channel, and (ii) cytoplasmic outlet pores sprinkle Na.sup.+ ions laterally over the inner surface of the plasma membrane to effect a rapid depolarization. VGSCs can be classified into two major categories. Category-I isoforms currently comprise nine highly homologous clones (Na.sub.v 1.1- Na.sub.v 1.9), most of which have been functionally expressed. In contrast, the category-II isoform consists of one clone (Na.sub.x), which has not been successfully expressed in an exogenous system. It is considerably different from the category-I isoforms, especially in the S4 segment, and shows little voltage dependence. The main function of the category-I isoforms is to form an action potential upstroke. However, Na.sub.v 1.6 can also influence subthreshold electrical activity in neurons through the "persistent" and "resurgent" Na.sup.+ currents, indicating that the VGSC itself can modulate overall neuronal firing behavior. Na.sub.v 1.8 and Na.sub.v 1.9 are preferentially expressed in peripheral nociceptive neurons and contain a structure common to tetrodotoxin (TTX)-resistant Na.sup.+ channels. Both Na.sub.v 1.8 and Na.sub.v 1.9 play a pivotal role in pain sensation. The category-II isoform Na.sub.x (x = unknown function) is a "concentration-sensitive" but not "voltage-sensitive" Na.sup.+ channel. It is involved in regulation of salt intake behavior by sensing an increase in [Na.sup.+]o, and it should be renamed as Na.sub.c (c = concentration).

ANSWER 7 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN L55

DUPLICATE

2002:35174982 BIOTECHNO ACCESSION NUMBER:

Specific contribution of human T-type calcium channel TITLE:

isotypes ($\alpha.sub.1.sub.G$, $\alpha.sub.1.sub.H$ and α.sub.1.sub.1) to neuronal excitability

Chemin J.; Monteil A.; Perez-Reyes E.; Bourinet E.; AUTHOR:

Nargeot J.; Lory P.

P. Lory, Inst. de Genetiq. Humaine, CNRS UPR 1142, 141 CORPORATE SOURCE:

rue de la Cardonille, F-34396 Montpellier Cedex 05,

France.

E-mail: philippe.lory@igh.cnrs.fr

Journal of Physiology, (01 APR 2002), 540/1 (3-14), 51 SOURCE:

reference(s)

CODEN: JPHYA7 ISSN: 0022-3751

Journal; Article DOCUMENT TYPE: COUNTRY: United Kingdom

LANGUAGE: English SUMMARY LANGUAGE: English 2002:35174982 BIOTECHNO

In several types of neurons, firing is an intrinsic property produced by AB specific classes of ion channels. Low-voltage-activated T-type calcium channels (T-channels), which activate with small membrane depolarizations, can generate burst firing and pacemaker activity. Here we have investigated the specific contribution to neuronal excitability of cloned human T-channel subunits. Using HEK-293 cells transiently transfected with the human $\alpha.sub.1.sub.G$ (Ca.sub.v3.1), $\alpha.sub.1.sub.H$ (Ca.sub.v3.2) and $\alpha.sub.1.sub.I$ (Ca.sub.v3.3) subunits, we describe significant differences among these isotypes in their biophysical properties, which are highlighted in action potential clamp studies. Firing activities occurring in cerebellar Purkinje neurons and in thalamocortical relay neurons used as voltage clamp waveforms revealed that $\alpha.sub.1.sub.G$ channels and, to a lesser extent, $\alpha.sub.1.sub.H$ channels produced large and transient currents, while currents related to $\alpha.sub.1.sub.G$ channels exhibited facilitation and produced a sustained calcium entry associated with the depolarizing after-potential interval. Using simulations of reticular and relay thalamic neuron activities, we show that $\alpha.sub.1.sub.I$ currents contributed to sustained electrical activities, while $\alpha.sub.1.sub.G$ and $\alpha.sub.1.sub.H$ currents generated short burst firing. Modelling experiments with the NEURON model further revealed that the $\alpha.sub.1.sub.G$ channel and $\alpha.sub.1.sub.I$ channel parameters best accounted for T-channel activities described in thalamocortical relay neurons and in reticular neurons, respectively. Altogether, the data provide evidence for a role of $\alpha.sub.1.sub.I$ channel in pacemaker activity and further demonstrate that each T-channel pore-forming subunit displays specific gating properties that account for its unique contribution to neuronal firing.

ANSWER 8 OF 19 PASCAL COPYRIGHT 2005 INIST-CNRS. ALL RIGHTS RESERVED. L55on STN

ACCESSION NUMBER: 2002-0416439 PASCAL

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reserved.

TITLE (IN ENGLISH): Block of the background K.sup.+ channel TASK-1

contributes to arrhythmogenic effects of

platelet-activating factor

BARBUTI Andrea; ISHII Satoshi; SHIMIZU Takao; ROBINSON AUTHOR:

Richard B.; FEINMARK Steven J.

CORPORATE SOURCE: Center for Molecular Therapeutics, Department of

Pharmacology, Columbia University, New York, New York 10032, United States; Department of Biochemistry and

Molecular Biology, University of Tokyo, Tokyo,

113-003, Japan

American journal of physiology. Heart and circulatory SOURCE:

physiology, (2002), 51(6), H2024-H2030, 26 refs.

ISSN: 0363-6135 CODEN: AJPPDI

DOCUMENT TYPE:

Journal Analytic BIBLIOGRAPHIC LEVEL:

United States COUNTRY:

LANGUAGE: English

INIST-670D, 354000100717110100 AVAILABILITY:

AN 2002-0416439 PASCAL Copyright .COPYRGT. 2002 INIST-CNRS. All rights reserved. Platelet-activating factor (PAF), an inflammatory phospholipid, induces ventricular arrhythmia via an unknown ionic mechanism. We can now link PAF-mediated cardiac electrophysiological effects to inhibition of a twopore domain K.sup.+ channel [TWIK-related acid-sensitive K.sup.+ background channel (TASK-1)]. Superfusion of carbamyl-PAF (C-PAF), a stable analog of PAF, over murine ventricular myocytes causes abnormal automaticity, plateau phase arrest of the action potential, and early afterdepolarizations in paced and quiescent cells from wild-type but not PAF receptor knockout mice. C-PAF-dependent currents are insensitive to Cs.sup.+ and are outwardly rectifying with biophysical properties consistent with a K.sup.+-selective channel. The current is blocked by TASK-1 inhibitors, including protons, Ba.sup.2.sup.+, Zn.sup.2.sup.+, and methanandamide, a stable analog of the endogenous lipid ligand of cannabanoid receptors. In addition, when TASK-1 is expressed in CHO cells that express an endogenous PAF receptor, superfusion of C-PAF decreases the expressed current. Like C-PAF, methanandamide evoked spontaneous activity in quiescent myocytes. C-PAF- and methanandamide-sensitive currents are blocked by a specific protein kinase C (PKC) inhibitor, implying overlapping signaling pathways. In conclusion, C-PAF blocks TASK-1 or a closely related channel, the effect is PKC dependent, and the inhibition alters the electrical activity of myocytes in ways that would be arrhythmogenic in the intact heart.

L55 ANSWER 9 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN

ACCESSION NUMBER:

2001:34073220 BIOTECHNO

TITLE: AUTHOR:

CP

AΒ

Electroporation in a model of cardiac defibrillation Ashihara T.; Yao T.; Namba T.; Ito M.; Ikeda T.;

Kawase A.; Toda S.; Suzuki T.; Inagaki A.; Sugimachi

M.; Kinoshita M.; Nakazawa K.

CORPORATE SOURCE:

Dr. T. Ashihara, First Dept. of Internal Medicine,

Shiga University of Medical Science, Seta Tsukinowa-cho, Otsu-city 520-2192, Japan.

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SOURCE:

AB

Journal of Cardiovascular Electrophysiology, (2001),

12/12 (1393-1403), 47 reference(s) CODEN: JCELE2 ISSN: 1045-3873

DOCUMENT TYPE:

Journal; Article

COUNTRY:

United States

LANGUAGE:

English

SUMMARY LANGUAGE:

English

AN 2001:34073220

BIOTECHNO

Introduction: It is known that high-strength shock disrupts the lipid matrix of the myocardial cell membrane and forms reversible aqueous pores across the membrane. This process is known as "electroporation." However, it remains unclear whether electroporation contributes to the mechanism of ventricular defibrillation. The aim of this computer simulation study was to examine the possible role of electroporation in the success of defibrillation shock. Methods and Results: Using a modified Luo-Rudy-1 model, we simulated two-dimensional myocardial tissue with a homogeneous bidomain nature and unequal anisotropy ratios. Spiral waves were induced by the S1-S2 method. Next, monophasic defibrillation shocks were delivered externally via two line electrodes. For nonelectroporating tissue, termination of ongoing fibrillation succeeded; however, new spiral waves were initiated, even with high-strength shock (24 V/cm). For electroporating tissue, high-strength shock (24 V/cm) was sufficient to extinguish ongoing fibrillation and did not initiate any new spiral waves. Weak shock (16 to 20 V/cm) also extinguished ongoing fibrillation; however, in contrast to the high-strength shock, new spiral waves were initiated. Success in defibrillation depended on the occurrence of electroporation-mediated anodal-break excitation from the physical anode and the virtual anode. Some excitation wavefronts following electrical shock used a

deexcited area with recovered excitability as a pass-through point; therefore, electroporation-mediated anodal-break excitation is necessary to block out the pass-through point, resulting in successful defibrillation. Conclusion: The electroporation-mediated anodal-break excitation mechanism may play an important role in electrical defibrillation.

ANSWER 10 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN L55 DUPLICATE

ACCESSION NUMBER: 1999:29243164 BIOTECHNO

Structure and function of cardiac potassium channels TITLE:

AUTHOR: Snyders D.J.

D.J. Snyders, Department of Molecular Biophysics, CORPORATE SOURCE:

Department of Biochemistry (UIA), University of

Antwerp, Universiteitsplein 1 - T4.21, B-2610 Antwerp,

Belgium.

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Cardiovascular Research, (1999), 42/2 (377-390), 114

reference(s)

CODEN: CVREAU ISSN: 0008-6363

PUBLISHER ITEM IDENT.:

SOURCE:

S0008636399000711

DOCUMENT TYPE: Journal; General Review

COUNTRY: Netherlands LANGUAGE: English

English SUMMARY LANGUAGE: 1999:29243164 **BIOTECHNO**

Recent advances in molecular biology have had a major impact on our understanding of the biophysical and molecular properties of ion channels. This review is focused on cardiac potassium channels which, in general, serve to control and limit cardiac excitability. Approximately 60 K.sup.+ channel subunits have been cloned to date. The (evolutionary) oldest potassium channel subunits consist of two transmembrane (Tm) segments with an intervening pore-loop (P). Channels formed by four 2Tm-1P subunits generally function as inwardly rectifying K.sup.+-selective channels (KirX.Y): they conduct substantial current near the resting potential but carry little or no current at depolarized potentials. The inward rectifier I(K1) and the ligand-gated K(ATP) and K(ACh) channels are composed of such subunits. The second major class of K.sup.+ channel subunits consists of six transmembrane segments (S1- S6). The S5-P-S6 section resembles the 2Tm-1P subunit, and the additional membrane-spanning segments (especially the charged S4 segment) endow these 6Tm-1P channels with voltage-dependent gating. For both major families, four subunits assemble into a homo- or heterotetrameric channel, subject to specific subunit-subunit interactions. The 6Tm-1P channels are closed at the resting potential, but activate at different rates upon depolarization to carry sustained or transient outward currents (the latter due to inactivation by different mechanisms). Cardiac cells typically display at least one transient outward current and several delayed rectifiers to control the duration of the action potential. The molecular basis for each of these currents is formed by subunits that belong to different Kvx.y subfamilies and alternative splicing can contribute further to the diversity in native cells. These subunits display distinct pharmacological properties and drug-binding sites have been identified. Additional subunits have evolved by concatenation of two 2Tm-1P subunits (4Tm-2P); dimers of such subunits yield voltage- independent leak channels. A special class of 6Tm-1P subunits encodes the 'funny' pacemaker current which activates upon hyperpolarization and carries both Na.sup.+ and K.sup.+ ions. The regional heterogeneity of K.sup.+ currents and action potential duration is explained by the heterogeneity of subunit expression, and significant changes in expression occur in cardiac disease, most frequently a reduction. This electrical remodelling may also be important for novel antiarrhythmic therapeutic strategies. The recent crystallization of a

2Tm-1P channel enhances the outlook for more refined molecular approaches.

ANSWER 11 OF 19 PASCAL COPYRIGHT 2005 INIST-CNRS. ALL RIGHTS RESERVED. L55

on STN

SOURCE:

AUTHOR:

ACCESSION NUMBER: 1998-0450199 PASCAL

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reserved.

Functional knockout of the transient outward current, TITLE (IN ENGLISH):

long-QT syndrome, and cardiac remodeling in mice

expressing a dominant-negative Kv4 α subunit

BARRY D. M.; HAODONG XU; SCHUESSLER R. B.; NERBONNE J. **AUTHOR:**

Department of Molecular Biology and Pharmacology, CORPORATE SOURCE:

> Washington University Medical School, St Louis, Mo, United States; Department of Surgery, Washington

University Medical School, St Louis, Mo, United States Circulation research, (1998), 83(5), 560-567, 34 refs.

ISSN: 0009-7330 CODEN: CIRUAL

Journal; Short communication DOCUMENT TYPE:

Analytic BIBLIOGRAPHIC LEVEL: COUNTRY: United States

LANGUAGE: English

INIST-7216, 354000070299530110 AVAILABILITY:

1998-0450199 PASCAL AN

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A novel in vivo experimental strategy, involving cell AB type-specific expression of a dominant-negative K.sup.+ channel

pore-forming α subunit, was developed and exploited to

probe the molecular identity of the cardiac transient outward K.sup.+ current (I.sub.t.sub.o). A point mutation (W to F) was introduced at

position 362 in the pore region of Kv4.2 to produce a

nonconducting mutant (Kv4.2W362F) subunit. Coexpression of Kv4.2W362F with Kv4.2 (or Kv4.3) attenuates the wild-type currents, and the effect is subfamily specific; ie, Kv4.2W362F does not affect heterologously expressed Kv1.4 currents. With the use of the α -myosin heavy chain promoter to direct cardiac-specific expression, several lines of Kv4.2W362F transgenic mice were generated. Electrophysiological recordings reveal that I.sub.t.sub.o is selectively eliminated in

ventricular myocytes isolated from transgenic mice expressing Kv4.2W362F,

thereby demonstrating directly that the Kv 4 subfamily underlies I.sub.t.sub.o in the mammalian heart. Functional knockout of

I.sub.t.sub.o leads to marked increases in action

potential durations in ventricular myocytes and to prolongation of the QT interval in surface ECG recordings. In addition, a novel rapidly activating and inactivating K.sup.+ current, which is not detectable in myocytes from nontransgenic littermates, is evident in

Kv4.2W362F-expressing ventricular cells. Importantly, these results demonstrate that electrical remodeling occurs in the

heart when the expression of endogenous K.sup.+ channels is altered.

ANSWER 12 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN

BIOTECHNO ACCESSION NUMBER: 1997:27289761

Molecular mechanism and functional significance of the TITLE:

> MinK control of the KvLQT1 channel activity Romey G.; Attali B.; Chouabe C.; Abitbol I.; Guillemare E.; Barhanin J.; Lazdunski M.

M. Lazdunski, Inst. Pharmacol. Molec./Cellulaire, CORPORATE SOURCE:

CNRS, 660 route des Lucioles, Sophia Antipolis, 06560

Valbonne, France.

E-mail: ipmc@unice.fr

Journal of Biological Chemistry, (1997), 272/27 SOURCE:

> (16713-16716), 28 reference(s) CODEN: JBCHA3 ISSN: 0021-9258

DOCUMENT TYPE:

Journal; Article

COUNTRY: LANGUAGE: United States

SUMMARY LANGUAGE:

English English

AN

1997:27289761 BIOTECHNO

The very slowly activating delayed rectifier K.sup. - channel I(Ks) is AB essential for controlling the repolarization phase of cardiac action potentials and K.sup. - homeostasis in the inner ear. The I(Ks) channel is formed via the assembly of two transmembrane proteins, KvLQT1 and MinK. Mutations in KvLQT1 are associated with a long QT syndrome that causes syncope and sudden death and also with deafness. Here, we show a new mode of association between ion channel forming subunits in that the cytoplasmic C-terminal end of Mink interacts directly with the pore region of KvLQT1. This interaction reduces KvLQT1 channel conductance from 7.6 to 0.58 picosiemens. However, because Mink also reveals a large number of previously silent KvLQT1 channels (x 60), the overall effect is a large increase (x 4) in the macroscopic K.sup.- current. Conformational changes associated with the KvLQT1/MinK association create very slow and complex activation kinetics without much alteration in the deactivation process. Changes induced by Mink have an essential regulatory role in the development of this K.sup.channel activity upon repetitive electrical stimulation with a particular interest in tachycardia.

ANSWER 13 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN L55 DUPLICATE

ACCESSION NUMBER:

BIOTECHNO 1996:26424228

TITLE:

Absence of the β subunit (cchb1) of the skeletal muscle dihydropyridine receptor alters expression of

the $\alpha.sub.1$ subunit and eliminates excitation-contraction coupling

AUTHOR:

Gregg R.G.; Messing A.; Strube C.; Beurg M.; Moss R.;

Behan M.; Sukhareva M.; Haynes S.; Powell J.A.;

Coronado R.; Powers P.A.

CORPORATE SOURCE:

R.G. Gregg, Waisman Center, University of Wisconsin,

1500 Highland Avenue, Madison, WI 53705, United

States.

SOURCE:

Proceedings of the National Academy of Sciences of the United States of America, (1996), 93/24 (13961-13966)

CODEN: PNASA6 ISSN: 0027-8424

DOCUMENT TYPE:

Journal: Conference Article

COUNTRY:

United States

LANGUAGE:

English English

SUMMARY LANGUAGE:

BIOTECHNO

AN 1996:26424228 The multisubunit $(\alpha(1S), \alpha.sub.2/\delta, \beta.sub.1, and$ AB γ) skeletal muscle dihydropyridine receptor transduces transverse tubule membrane depolarization into release of Ca.sup.2.sup.+ from the sarcoplasmic reticulum, and also acts as an L-type Ca.sup.2.sup.+ channel. The $\alpha(1S)$ subunit contains the voltage sensor and channel pore, the kinetics of which are modified by the other subunits. To determine the role of the β .sub.1 subunit in channel activity and excitation-contraction coupling we have used gene targeting to inactivate the $\beta.sub.1$ gene. $\beta.sub.1$ -null mice die at birth from asphyxia. Electrical stimulation of β .sub.1-null muscle fails to induce twitches, however, contractures are induced by caffeine. In isolated β .sub.1-null myotubes, action potentials are normal, but fail to elicit a Ca.sup.2.sup.+ transient. L-type Ca.sup.2.sup.+ current is decreased 10- to 20-fold in the β .sub.1null cells compared with littermate controls. Immunohistochemistry of cultured myotubes shows that not only is the $\beta.\,sub.\,1$ subunit absent, but the amount of $\alpha\,(1S)$ in the membrane also is undetectable. In contrast, the β .sub.1 subunit is

localized appropriately in dysgenic, mdg/mdg, ($\alpha(1S)$ -null)

cells. Therefore, the β .sub.1 subunit may not only play an important role in the transport/insertion of the $\alpha(1S)$ subunit into the membrane, but may be vital for the targeting of the muscle dihydropyridine receptor complex to the transverse tubule/sarcoplasmic reticulum junction.

ANSWER 14 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN L55

DUPLICATE

ACCESSION NUMBER:

1996:26266146 BIOTECHNO

Class III antiarrhythmic effects of zatebradine: TITLE:

Time-, state-, use-, and voltage-dependent block of

hKv1.5 channels

Valenzuela C.; Delpon E.; Franqueza L.; Gay P.; Perez AUTHOR:

O.; Tamargo J.; Snyders D.J.

Inst. of Pharmacology and Toxicology, School of CORPORATE SOURCE:

Medicine, Universidad Complutense, 28040 Madrid, Spain.

Circulation, (1996), 94/3 (562-570) SOURCE:

CODEN: CIRCAZ ISSN: 0009-7322

DOCUMENT TYPE:

Journal; Article United States

COUNTRY: LANGUAGE:

English

SUMMARY LANGUAGE:

English

AN 1996:26266146 BIOTECHNO

Background: Zatebradine is a bradycardic agent that inhibits the AΒ hyperpolarization-activated current (I(f)) in the rabbit sinoatrial node. It also prolongs action potential duration in papillary muscles in guinea pigs and in Purkinje fibers in rabbits. The underlying mechanism by which zatebradine induces this effect has not been explored, but it is likely to involve K.sup.+ channel block. Methods and Results: Cloned human cardiac K.sup.+ delayed rectifier currents (hKv1.5) were recorded in Ltk cells transfected with their coding sequence. Zatebradine 10 μ mol/L did not modify the initial activation time course of the current but induced a subsequent decline to a lower steady-state current level with a time constant of 109±16 ms. Zatebradine inhibited hKv1.5 with an apparent K(D) of 1.86±0.14 µmol/L. Block was voltage dependent (electrical distance $\delta \text{=0.177} \pm \text{0.003})$ and accumulated in a use-dependent manner during 0.5- and 1-Hz pulse trains because of slower recovery kinetics in the presence of the drug. Zatebradine reduced the tail current amplitude, recorded at -30 mV, and slowed the deactivation time course, which resulted in a 'crossover' phenomenon when control and zatebradine tail currents were superimposed. Conclusions: These results indicate that (1) zatebradine is an open-channel blocker of hKv1.5, (2) binding occurs in the internal mouth of the ion pore, (3) unbinding is required before the channel can close, and (4) zatebradine-induced block is use dependent because of slower recovery kinetics in the presence of the drug. These effects may explain the prolongation of the cardiac action potential and could be clinically relevant.

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95:116217 LIFESCI ACCESSION NUMBER:

Surfing the DNA databases for K super(+) channels nets yet TITLE:

more diversity

Salkoff, L.; Jegla, T. AUTHOR:

Dep. Anat. and Neurobiol., Washington Univ. Sch. Med., St. CORPORATE SOURCE:

Louis, MO 63110, USA

NEURON, (1995) vol. 15, no. 3, pp. 489-493. SOURCE:

ISSN: 0896-6273.

DOCUMENT TYPE: Journal FILE SEGMENT: LANGUAGE: English

K super(+) channels are a life and death matter. Perhaps the best assessment of whether a cell is living or dead is whether or not

it has a membrane potential, and K super(+) channels have a significant

role in setting the membrane potential in cells from a wide variety of life forms. In addition to this life or death matter, K super(+) channels serve a host of other functions relating to the electrical lives of cells, like setting the frequency and duration of action potentials and, in general, shaping the electrical activity of cells. Because multiple K super(+) channel types have also been found in a wide variety of cells that are not known to be electrically excitable, it is likely that we don't yet comprehend all of the functions of these versatile proteins. Data now surfacing from the various genomic DNA sequencing projects suggest that this family of proteins might be even more diverse than previously imagined. The TOK1 channel is novel, not only because of its unique subunit structure (it resembles two K super(+) channel subunits of different classes linked together) and its unique physiology (it is outwardly rectifying by a nonconventional mechanism), but also because of the way in which this unique channel was found. Rather than using the conventional experimental methods of molecular biology in a wet lab, the authors discovered TOK1 by surfing the public DNA database for one of the rare and special ancient conserved protein regions that have been identified as a result of the genome sequencing projects. This particular ancient conserved region is the universal signature of the ion-selective pore of K super(+) channels.

ANSWER 16 OF 19 BIOTECHNO COPYRIGHT 2005 Elsevier Science B.V. on STN

DUPLICATE

ACCESSION NUMBER:

1993:23197520 BIOTECHNO

TITLE:

The cloning and expression of a sodium channel

β1-subunit cDNA from human brain

AUTHOR:

McClatchey A.I.; Cannon S.C.; Slaugenhaupt S.A.;

Gusella J.F.

CORPORATE SOURCE:

Molecular Neurogenetics Unit, Massachusetts General Hospital East, 13th Street, Charlestown, MA 02129,

United States.

SOURCE:

Human Molecular Genetics, (1993), 2/6 (745-749)

CODEN: HMGEE5 ISSN: 0964-6906

DOCUMENT TYPE:

Journal; Article

COUNTRY: LANGUAGE: United Kingdom

English

SUMMARY LANGUAGE:

English

1993:23197520 BIOTECHNO

Electrical excitability of neurons and muscle cells AB is mediated largely through the actions of the voltage-gated sodium channel. Initiation and propagation of the action potential is a direct result of the precisely controlled inward flux of sodium through these channels. Much attention has been paid to the sodium channel α -subunit, the major, pore-forming component. However, α -subunits are associated with one or more smaller β -subunits, which have been implicated in the critical fine tuning of the gating properties of the channel. To investigate the properties of the β -subunit, we have isolated a cDNA encoding the human brain β 1-subunit and assigned the corresponding gene to chromosome 19. We have also examined the effects of expressing the brain β1-subunit on the kinetics of a coexpressed muscle sodium channel $\alpha\mbox{-subunit.}$ Our results underscore the functional importance of the β 1-subunit and imply a conserved mechanism for the interaction of the $\beta1$ -subunit with different α -subunits.

ANSWER 17 OF 19 PASCAL COPYRIGHT 2005 INIST-CNRS. ALL RIGHTS RESERVED. L55

on STN

ACCESSION NUMBER: 1993-0523472

TITLE (IN ENGLISH):

The cloning and expression of a sodium channel

β1-subunit cDNA from human brain

PASCAL

AUTHOR:

MCCLATCHEY A. I.; CANNON S. C.; SLAUGENHAUPT S. A.;

GUSELLA J. F.

CORPORATE SOURCE: Massachusetts gen. hosp., molecular neurogenetics

unit, Charlestown MA 02129, United States

SOURCE: Human molecular genetics, (1993), 2(6), 745-749, 27

refs.

DOCUMENT TYPE: Journal BIBLIOGRAPHIC LEVEL: Analytic

COUNTRY: United Kingdom

LANGUAGE: English

AVAILABILITY: INIST-22540, 354000033976850210

AN 1993-0523472 PASCAL

Blectrical excitability of neurons and muscle cells is mediated largely through the actions of the voltage-gated sodium channel. Initiation and propagation of the action potential is a direct result of the precisely controlled inward flux of sodium through these channels. Much attention has been paid to the sodium channel α -subunit, the major, pore-forming component. However, α -subunits are associated with one or more smaller β -subunits, which have been implicated in the critical fine tuning of the gating properties of the channel. To investigate the properties of the β -subunit, we have isolated a cDNA encoding the human brain β 1-subunit and assigned the corresponding gene to chromosome 19. We have also examined the effects of expressing the brain

β1-subunit on the kinetics of a coexpressed muscle sodium channel

L55 ANSWER 18 OF 19 LIFESCI COPYRIGHT 2005 CSA on STN

ACCESSION NUMBER:

α-subunit

91:70123 LIFESCI

TITLE:

Noninvasive recording of receptor cell action potentials and sustained currents

from single taste buds maintained in the tongue: The

response to mucosal NaCl and amiloride.

AUTHOR: Avenet, P.; Lindemann, B.

CORPORATE SOURCE: Dep. Physiol., Univ. Saarlandes, D-6650 Homburg/Saar, FRG

SOURCE: J. MEMBR. BIOL., (1991) vol. 124, no. 1, pp. 33-41.

DOCUMENT TYPE: Journal FILE SEGMENT: M; R LANGUAGE: English SUMMARY LANGUAGE: English

AB Apical membrane currents were recorded from the taste pore of single taste buds maintained in the tongue of the rat, using a novel approach. Under a dissection microscope, the 150- mu m opening of a saline-filled glass pipette was positioned onto single fungiform papillae, while the mucosal surface outside the pipette was kept dry.

Electrical responses of receptor cells to chemical stimuli, delivered from the pipette, were recorded through the pipette while the cells remained undamaged in their natural environment. We observed monophasic transient currents of 10-msec duration and 10-100 pA amplitude, apparently driven by action potentials arising spontaneously in the receptor cells.

L55 ANSWER 19 OF 19 LIFESCI COPYRIGHT 2005 CSA on STN

ACCESSION NUMBER: 82:65756 LIFESCI

TITLE: Ionic channels in skeletal muscle.
AUTHOR: Stefani, E.; Chiarandini, D.J.

CORPORATE SOURCE: Dep. Physiol., Cent. Estudios Avanzados del Inst.

Politecnico Nacil., Apartado Postal 14-740, Mexico 14,

D.F., Mexico

SOURCE: ANNU. REV. PHYSIOL., (1982) vol. 44, pp. 357-372.

DOCUMENT TYPE: Journal

TREATMENT CODE: General Review

FILE SEGMENT: M

LANGUAGE: English

AB There is now considerable evidence that ions can move across the **cell** membrane through voltage-gated aqueous **pores** celled

"ionic channels." Each channel has a characteristic permeability, selectivity, and kinetics. **Electrical** excitation in skeletal muscle involves voltage- and time-dependent changes of the permeabilities to Na super(+) and K super(+) which induce a transient inflow of Na super(+) into the fiber followed by an outflow of K super(+). As a result of these ionic movements the **action potential** is generated. Besides these Na super(+) and K super(+) channels, a voltage-dependent Ca super(2+) channel has been described recently in frog skeletal muscle. At rest, Cl super(-) and other K super(+) channels are responsible for the dominant conductance. Most of these K super(+) channels rectify inward current. Here The authors review recent research on ionic channels in twitch and tonic skeletal widely used.